

**RESULTS OF INVESTIGATION:** Analysis showed that the article failed to meet the test specified in the National Formulary regarding permissible variations in the weight of individual containers. Analysis showed also that the individual containers contained from 42 percent to 117 percent of the declared amount of amobarbital.

**LIBELED:** 8-5-55, E. Dist. Va.

**CHARGE:** 501 (b)—the article, when shipped, purported to be and was represented as "Sodium Amobarbital," a drug the name of which is recognized in the National Formulary, an official compendium, and its strength differed from the standard set forth in such compendium; and 502 (a)—the label statement "Amobarbital Sodium 7½ gr." was false and misleading.

**DISPOSITION:** 12-6-55. Default—destruction.

**4935. Code #55 capsules.** (F. D. C. No. 38097. S. No. 19-715 M.)

**QUANTITY:** 2 cartons, 7,850 capsules each, at Columbus, Ohio.

**SHIPPED:** 1-11-52 and 11-13-52, from Detroit, Mich.

**RESULTS OF INVESTIGATION:** Analysis showed that the article contained less than the declared amount of vitamin C (ascorbic acid).

**LIBELED:** 7-20-55, S. Dist. Ohio.

**CHARGE:** 501 (c)—the strength of the article, while held for sale, differed from that which it purported and was represented to possess, namely, 50 milligrams of vitamin C per capsule; and 502 (a)—the label statement "Ingredients in each capsule: \* \* \* Ascorbic Acid U. S. P. 50 Mg." was false and misleading.

**DISPOSITION:** 8-25-55. Default—destruction.

**4936. Moe Pap liquid.** (F. D. C. No. 38101-A. S. No. 13-976 M.)

**QUANTITY:** 100 4-oz. btls. at Memphis, Tenn.

**SHIPPED:** 4-15-55 and 4-26-55, from St. Louis, Mo.

**RESULTS OF INVESTIGATION:** Analysis showed that the article contained less than 75 percent of the declared amount of vitamin B<sub>1</sub>.

**LIBELED:** 7-25-55, W. Dist. Tenn.

**CHARGE:** 501 (c)—the strength of the article, while held for sale, differed from that which it purported and was represented to possess, namely, 1,500 I. U. of vitamin B<sub>1</sub> per fluid ounce; and 502 (a)—the label statement "Thiamine Hydrochloride (Vitamin B<sub>1</sub>) 1500 I. U. per Fluid Ounce" was false and misleading.

**DISPOSITION:** 9-1-55. Default—destruction.

**4937. Ala-Dyne tablets.** (F. D. C. No. 38238. S. No. 29-309 M.)

**QUANTITY:** 1 1,000-tablet btl. and 608 100-tablet btls. at Emerson, N. J., in possession of Allied Drugs, Inc.

**SHIPPED:** 12-19-50 and 5-26-52, from Cleveland, Ohio.

**LABEL IN PART:** (Btl.) "Ala-Dyne Each Tablet Contains Acetylsalicylic Acid 4 grs. Calcium Glutamate 2 grs. Ascorbic Acid 30 mg. Allied Drugs, Inc. Hackensack, New Jersey Distributors Caution To be dispensed by or on the prescription of a physician. 2676 [or "4993"]."

**RESULTS OF INVESTIGATION:** The article was shipped in interstate commerce in bulk, and, upon receipt by the consignee, was repackaged. Analysis showed that lot number 2676 contained 76 percent of the labeled amount of acetylsalicylic acid.